

Form P 1

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APPLICATION FOR A PATENT AND ACKNOWLEDGEMENT OF RECEIPT [Section 30(1) - Regulation 22]

Offi	cial Ap	pplication No	Applicants Ref	7
21	10	76 2382	64942	
71		Full names of Applicant		
	JG PI	LUS INTERNATIONAL INC.		
		Address of Applicant		
5	50,	Chambly, Sherbrooke, Quebec	JlJ 2Y2, CANADA	
54		Title of invention		
D	RY C	CHLORINATION OF PGM-BEARING C	HROMITE ORES	

The application is accompanied by :-

✓	1.	Two copies of a complete specification of 22 pages
✓	2.	Informal Drawings of one sheet
1	3.	Publication Particulars and abstract
	4.	Λ copy of Figure 1 of the drawings for abstract
	5.	Assignment of invention
	6.	Certified priority documents
	7.	Translation of priority documents
	8.	An assignment of priority rights
	9.	A copy of Form P2 and the specification of S A
		Patent Application No 21. 01.
	10	A declaration and power of attorney on Form P3
	11	Request for ante-dating on Form P 4
	12	Request for classification on Form P9
1	13	Form P2
		Address for Service John Galgut Johannesburg

Dated this 26 March 1996

Applicant's Patent Attorney

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PRETORIA

REGISTRATEUR VAN PATENTE, MODFLLE, HANDELSMERKE EN DUTEURSREG

REPUBLIC OF SOUTH AFRICA PATENTS ACT, 1978 COMPLETE SPECIFICATION

[Section 30(1) - Regulation 28]

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Full names of Applicant(s)	
71 UG PLUS INTERNATIONAL INC.	
Full parada as damentas (a)	•
Full name(s) of inventor(s)	

Tit	le of invention
54	DRY CHLORINATION OF PGM-BEARING CHROMITE ORES

Jean-Marc LALANCETTE and Mario BERGERON

TITLE OF THE INVENTION

Dry chlorination of PGM-bearing chromite ores.

FIELD OF THE INVENTION

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The present invention pertains to a process for recovering PGM from PGM-bearing chromite ores.

BACKGROUND OF THE INVENTION

The term "Platinum Group Metals" (PGM) designates the six platinoids, namely, ruthenium, rhodium, palladium, osmium, iridium and platinum. These metals are often found together in variable proportions, and platinum normally predominates.

PGM are generally associated with the presence of ultramafic rocks containing agents such as nickel or chromium compounds, either sulfides or oxides. In fact, chromite ($Fe0 \cdot Cr_20_3$) has been observed to be associated with sulfides of the PGM in some important geological formations, such as the Bushveld Complex found in South Africa (Merensky Reef and UG2 Chromitite).

The current technology used for the recovery of PGM from these chromites ores calls for the grinding of the ore to about 75 microns (-200 mesh) followed in certain cases by gravity concentration and then by flotation. The concentrate of the sulfides containing the PGM and other sulfides, such as iron copper and nickel sulfides, is then smelted so as to produce a matterich in PGM. The acid digestion of the matte leads to the individual components of the PGM after rather elaborate chemical operations.

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The froth flotation process is a technique of general use in sulfides processing. It allows the production of base metal concentrates from ores with relatively low sulfides content. Copper and nickel productions, for example, depend upon enrichment by flotation to achieve high yield from ores having sulfide contents at the level of a few percents.

In the case of the PGM, the froth collected from the flotation of the chromite ores has a PGM content of 100 to 400 ppm and incorporates copper sulfide and nickel sulfide at the percent level.

Because of a number of factors, the flotation of the sulfides from chromite ore is rather delicate and much influenced by variations in ore composition induced by oxidation, occlusion in chromite (coarse grinding) and other variables responsible for the modifications of the surface of the particles. Although nominal recovery of the order of 80% are reported for the flotation of the PGM and associated sulfides from chromite ores, the actual practice can lead to significantly lower figures, and constant adjustment of the operational parameters are required. This situation and the fact that some types of PGM-bearing minerals cannot be floated readily makes desirable the investigation of alternate methods for the recovery of PGM from chromite ores.

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The analytical procedures for the determination of PGM as generally practiced call for the oxidation of the substrate in the presence of aqua regia, a mixture of hydrochloric and nitric acids, the presence of a free halogen, such as chlorine, being required for the more refractory elements of the group (see J.C. Van Loon and R.R. Barefoot, Determination of the Precious Metals, pp. 55,101, John Wiley & sons, 1991). Another approach calls

for a fusion with nickel sulfide followed by acid digestion of the melt under oxidizing conditions. Such procedures are deemed fairly efficient in converting all of the PGM in the form of soluble complexed chlorides that can be determined in solution by instrumental analysis.

although appropriate for analytical purposes, cannot be retained for industrial production. The amount of chemicals required for the total digestion of the PGM ore would render the cost of the operation absolutely prohibitive. Chromite ores contains typically from 20 to 30 percent chromite (Fe0 • Cr203) associated with 3-4 ppm of PGM. Since the standard oxidizing agents cited above are reported as strong enough to oxidize readily the iron oxide and trivalent chromium oxide of the chromite, the consumption of oxidizer per unit weight of PGM would become so high as to remove any profitability from the process.

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The examination of the chemical literature concerning the oxidation of ferrous oxide or trivalent

chromium oxide is quite convincing to the effect that oxidizing agents strong enough to oxidize PGM will very readily oxidize ferrous oxide or trivalent chromium oxide to ferric oxide and hexavalent chromium oxide.

Ferrous oxide is readily oxidized by air in the presence of acids such as nitric acid or hydrochloric acid (see S.M. Latimer, The oxidation state of the elements, 2nd ed., p. 224, Prentice-Hall, 1952). Even chromite itself, Fe0•Cr₂O₁, is reported oxidized by air, which is a much milder oxidizer than chlorine or nitric acid. This reaction is the industrial method of preparation of sodium chromate:

2 Fe0 • Cr₂0, + 4Na₂CO, + 3.50, - 4Na₂CrO, + 4CO, + Fe₂O,

(see H. Remy, Treatise on Inorganic Chemistry, Vol. II,
p. 155, Elsevier Publishing Company, 1956).

STATEMENT OF THE INVENTION

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While attempting to purify PGM-bearing chromites from the base metal components, the inventors have found that, under appropriate conditions, chlorine could react with the PGM in chromites without significant attack on chromite itself. This result, unexpected from

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the statements of the chemical literature as indicated above, opens the way to a new approach to the recovery of PGM from chromite ores.

The present invention therefore provides, according to one aspect, a process for recovering PGM from PGM-bearing chromite ores comprising the steps of:

- dry chlorinating said ores, conveniently at temperatures from 350°C to 800°C, to produce chlorinated ores;
- digesting said chlorinated ores with hydrochloric acid in the presence of chlorine to produce soluble
 PGM salts;
 - filtering residual insoluble chromite ores to provide substantially unaltered chromites and a pregnant solution containing PGM and a small quantity of other metals; and
 - recovering PGM from said pregnant solution.

In one form of the invention, the dry chlorinating step is conducted in the presence of sodium chloride.

In another form of the invention, the ores are heated prior to the dry chlorinating step.

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The invention presents several advantages over the pyrometallurgical approach currently used for the recovery of PGM from chromite ores, namely:

flotation is not required (although a concentrate by still be used, if available, with the present invention).

This step, which relies on physical properties of the ore (surface characteristics and size of sulfide particles), calls for constant adjustments for the flotation, and some ores are not amendable to it;

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- the percentage of recovery of PGM is from 10 to 20% higher than what is obtained in practice with the pyrometallurgical approach;
- the osmium which is almost completely lost

 in the pyrometallurgical approach is largely recovered in
 the condenser of this process.
 - the circuit of the present invention is less elaborate than the pyrometallurgical circuit, no recycling of slag or chemical digestion of matte being required to reach the stage of dissolved PGM; and

- the present invention can be operated on a modular basis and is less capital incentive than the pyrometallurgical approach.

Other objects and further scope of applicability of the present invention will become apparent from the detailed description given hereinafter. It should be understood, however, that this detailed description, while indicating preferred embodiments of the invention, is given by way of illustration only, since various changes and modifications within the spirit and scope of the invention will become apparent to those skilled in the art.

IN THE DRAWINGS

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Figure 1 is a block diagram illustrating the
various steps of one embodiment of a dry chlorination of
chromite ores made in accordance with present invention.

DESCRIPTION OF PREFERRED EMBODIMENTS

First, the ore is reduced to an appropriate size. It will be readily understood that too coarse a

material will prevent access to the values inside the particles and will slow down the reaction. On the other hand, very fine grinding involves significant cost. A value of particles from 75 microns to 45 microns appears as a practical compromise between reactivity and cost of grinding.

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This finely ground material is then dried to less than 0.1% of free water and heated up to the appropriate temperature for the dry chlorine treatment. This temperature of dry chlorination has been found to be variable depending on the refractoriness of the ore under treatment. With certain ores, a temperature of 450°C has been sufficient to recover more than 90% of the PGM. In other instances, higher temperatures (up to 700°C) have been required to achieve similar recovery. The temperature of dry chlorination is an important factor in the capacity of treatment of a given apparatus. temperature must be high enough to ensure the desired reactions with the PGM, while too high a temperature may prove detrimental to the equipment and may induce undesirable reactions with chromite. If the variations of reactivity of the ores are taken into account, it has

been found that a range of temperature from 350°C to 800°C allows to obtain good reactivity, leading to high percentage of extraction of PGM while limiting to acceptable values the side reactions with chromite. The ground ore can be heated to the selected temperature prior to contacting with chlorine or heated up in the chlorinator itself. Both procedures have been tried but the preheating approach turned out to be simpler, a rotary kiln or a fluidized bed being favored.

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The hot ore is then introduced in the dry chlorinator, which is kept at the selected temperature, and a slow stream of chlorine is circulated through the mass at constant temperature. It must be noted that the amount of chlorine required is very small and amounts from 1.0 to 1.5 times the combined chlorine in the course of the dry chlorination. The amount of chlorine consumed by the PGM is relatively negligible, the consumption being related to the amount of reacting chromite and base metals, such as iron, nickel and copper present in the starting ore. As an indicative estimate, it can be said that some of the base metals (Cu, Ni) at most a few percents of the iron in the chromite and less than one-

tenth of one percent of the chromium in the chromite will be converted into the corresponding soluble chlorides. In order to ensure an homogeneous distribution of the chlorine through the reacting mass, the chlorine stream can be diluted by a carrier gas, such as nitrogen for The dry chlorination can be achieved on a continuous basis or batch-wise by simple percolation through the mass of the reacting gas or by the fluidization of the reacting bed or by renewal of the surfaces in a rotating kiln. The contacting time required is relatively long, of the order of half an hour to two hours, depending on the nature of the ore, the temperature used and, to some extent, on the granulometry of the ore. For sake of simplicity of an apparatus which operates under rather aggressive conditions (chlorine at 600°C for example), a static bed of appropriate thickness has been found convenient. Since the material to be treated is to be loaded hot, the heating of the dry chlorination reactor has to be just enough to compensate for heat losses of the system.

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The volatiles leaving the dry chlorinator are directed to a condensing tower and quenched by a spray of

hydrochloric acid at a concentration of 6M. The temperature in this tower is of the order of 50°C and the condensable are collected and carried down by the acid spray.

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The solid left in the dry chlorinator after treatment is dumped after cooling in a digester containing the acid from the spray in the condensing tower, and is further digested, at 100°C, for one to several hours, the stirring of the mass being ensured by recirculation of the off gases of the condensation tower through the slurry. The off gases from the digester, mostly nitrogen, hydrochloric acid vapors, chlorine and traces of sulfur compounds, are directed back to a gas treatment system after recycling condensed hydrochloric acid to the digester.

The temperature and duration of digestion in hydrochloric acid in the presence of chlorine must be adjusted in such a fashion as to ensure the complete solution of the PGM. The heat increases the rate of such reactions, which would proceed nevertheless at room temperature but at a slower rate.

For sake of completeness of dissolution as readily as possible of the PGM, hot treatment is preferred over cold treatment and a one to several hours of contacting is retained.

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The dry chlorination, followed by digestion, has now transformed more than 90% of the PGM into soluble chlorinated entities. The addition of some sodium chloride in the system has produced soluble chlorides with some of the PGM that would otherwise give insoluble chlorides. In the case of iridium, for example, this metal gives a trichloride with chlorine at 500°C, but this salt is not water-soluble. However, in the presence of sodium chloride, at 600°C, the action of chlorine on Ir leads to sodium hexachloroiridate Na₂IrCl₆, a soluble salt. A similar behavior is noted with rhodium and ruthenium (see J.C. Van Loon et al, cited above).

The pregnant solution is then separated from the insoluble chromite ore by filtration and the cake is submitted to several washings in order to minimize the retention of the valuable species. After the rinsing

operations, the acid concentration in the pregnant solution is of the order of 3 molar.

The recovery of valuable species from this solution can be achieved by a variety of techniques, such as precipitation over activated carbon, or circulation over ion-exchange resins. The resulting depleted solution is directed to a distillation line in order to recycle the hydrochloric acid at a concentration of 6M which corresponds to the azeotropic composition of the HCl-H₂O system. The bottoms of this distillation contain the iron, the chromium and the base metals (Ni, Cu, etc...) that have been dissolved in the course of the digestion of the dry-chlorinated ore. This material is directed to the gas treatment system which also receives the vapors of the digester.

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The entities found at the inlet of the purification system are oxidizing by nature (Cl₂) and acidic. They also include all the ions liberated by the digestion. This mixture is reduced by percolation over iron filling, and the acidic lixiviate is neutralized with limestone. Under alkaline conditions, base metal

oxides are precipitated as hydroxides; hexavalent chromium, after reduction, is precipitated by the relatively large amounts of trivalent iron present, and the chlorine ends up as calcium chloride. The resulting slurry can be filtered and calcium chloride eliminated as solid after saturation of the circuit with this salt.

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Referring to TABLE 1, six examples are given to illustrate the implementation of the invention. The PGMbearing chromite ore (100 g) is reduced to the appropriate size, heated up at an appropriate temperature with or without addition of salt, in the presence of chlorine, and then digested in 6M hydrochloric acid for a determined period of time. The reacting mass is then filtered and rinsed; the filtrate, after removal of excess hydrochloric acid, is contacted with either activated carbon or ion-exchange resins to recover PGM. The analyses are done by ICP-MS technique.

In the case of Example 2, the pregnant solution after filtration of the insolubles contained 30 ppm of chromium and 50,000 ppm of iron. The acid was fractionated by distillation and the bottoms treated by

a slight excess of limes so as to give a neutral filtrate and a solid cake where ferric hydroxide predominates.

Although the invention has been described above with respect with one specific form, it will be evident to a person skilled in the art that it may be modified and refined in various ways. It is therefore wished to have it understood that the present invention should not be limited in scope, except by the terms of the following claims.

TABLE I

Examples	Source of ore	Particle	Dry	Dry chlorination	lon	Dic	Digestion)d	***50
No	(chromite)	Size							
		(micron)	Temp.	Duration	NaCl	јешр.	Duration	Initial	Recovery
			(C)	(hr.)		(°)	(hz.)	(<u>യർർ</u>)	0/0
H	Coleraine	75	350	1.0	or Or	100	m	4.50	A R
	Canada								3
7	Coleraine	45	450	1.5	or	00:	u	A 5.0	30
	Canada						1))	0
en .	Eastern Platinum	75	350	1.0	ou	00:	7	7.05 4	C
	South Africa						•))
4	Eastern Platinum	75	450	1.0	ou	305	7	4 507	,,,
	South Africa	1					•)) ,	0
S	Eastern Platinum	45	099	(r-1	yes	001	3	4.507	0
	South Africa								? .
9	Eastern Platinum	4.5	670	1.5	yes	100	4	4.507	0.7
	South Africa				ı		-		`
* Total values	for Pt, Pd,	Rh, Ru, Ir.	o C						
			24.						

The embodiments of the invention in which an exclusive property or privilege is claimed are defined as follows:

- 5 1. A process for recovering PGM from PGMbearing chromite ores, comprising the steps of:
 - dry chlorinating said ores at temperatures from 350°C to 800°C to produce chlorinated ores;
- digesting said chlorinated ores with hydrochloric acid in the presence of chlorine to produce soluble PGM salts;
 - filtering residual insoluble chromite ores to provide substantially unaltered chromites and a pregnant solution containing PGM and a small quantity of other metals; and
 - recovering PGM from said pregnant solution.
- 2. A process as defined in claim 1, wherein
 the dry chlorinating step is conducted in the presence of sodium chloride.

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CLAIMS

- 1. A process for recovering PGM from PGM-bearing chromite ores comprising the steps of:
- dry chlorinating said ores to produce chlorinated ores;
- digesting said chlorinated ores with hydrochloric acid in the presence of chlorine to produce soluble PGM salts;
 - filtering residual insoluble chromite ores to provide substantially unaltered chromites and a pregnant solution containing PCM and a small quantity of other metals; and
 - recovering PGM from said pregnant solution.
 - 2. A process as defined in claim 1 wherein the step of dry chlorinating said ores is carried out at temperatures from 350°C to 800°C.
 - 3. A process as defined in claim 1 or 2, wherein the dry chlorinating step is conducted in the presence of sodium chloride.

- 4. A process as defined in claim 2, comprising the step of heating said ores prior to said dry chlorinating step.
- 5. A process as defined in claim 4 comprising the step of grinding said chromite ores to a size of about 45 microns before the heating step.

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- 6. A process as defined in any one of the preceding claims comprising the step of cooling said chlorinated ores prior to the digesting step.
- 7. A process as defined in claim 6 wherein said digesting step is carried out in the presence of heat.
 - 8. A process as defined in any one of the preceding claims, comprising the step of condensing volatiles created by the dry chlorinating step, said volatiles including volatile salts of PGM.
 - 9. A process as defined in claim 8 wherein the condensing step is conducted by quenching said volatiles with a spray of hydrochloric acid.

A process as defined in claim 9, wherein said condensing step is carried out at a temperature in the order of 50°C.

11. A process as defined in claim 9, further directing said spray of hydrochloric acid to the digesting step.

12. A process as defined in claim 11, wherein said digesting step is conducted in the presence .. heat in the order of 100°C.

13. A process as defined in any one of the preceding claims, wherein gases resulting from the digesting step are directed to a gas treatment system after recycling condensed hydrochloric acid to the digesting step.

15 14. A process as defined in any one of the preceding claims, wherein said recovering is carried out by precipitating PGM in said pregnant solution over activated carbon.

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15. A process as defined in any one of the preceding claims, wherein said recovering step is carried out by circulating said pregnant solution over ion-exchange resins.

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16. A process as defined in claim 14 or 15 further comprising the step of distillating a depleted solution resulting from the recovering step to thereby recycle the hydrochloric acid; the bottoms of said distillating step being directed to a system for treating vapours from the digesting step.

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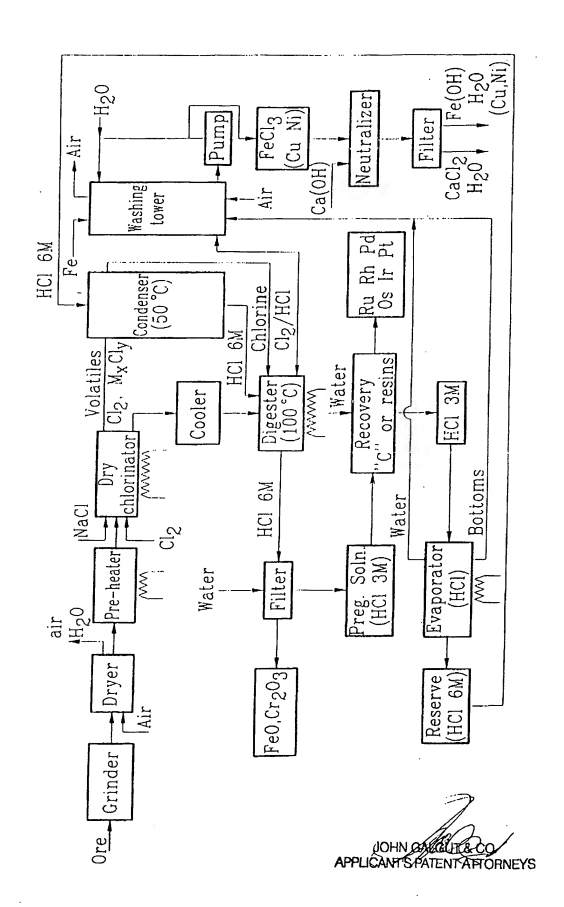
17. A process for recovering PGM from PCM bearing chromite ores substantially as hereinbefore described and as illustrated in Figure 1 of the accompanying drawings.

Original dated this 26th March, 1996

(Sgd) J Galgut JOHN GALGUT Applicant's Patent Attorney

Dated this 13 December 1996

Applicant's Patent Attorney



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